

Scapencing of 5 and of phaskIII

W Sequinase Kit

Purpose: Determine if phaskIII encodés a Known protein (e.g. CD36 or Limp: II)

Followed Kit instructions. Used primers: T7 - on plasmid pcDNAI 05A1.3 - in cDNA sequence 5' CTG TCG CTG TCC CCC TTC AG 3' short read

GGTACCGAGCTCGATCCACTAGTAACGGCCGCCAGTGTCTCTAAACGCCACCTGCAGGGCTACTG CTGCTCCGGCCACTGCCTGAGACTCACCTTGCTGGAACGTGAGCCTCGGCTTCTGTCATCTCTG

long read

ACTCACCTTGCTGGAACGTGAGCCTCGGCTTCTGTCATCTCTGTGGCCTCTGTCGCTTCTGTCGCTGTCCCCTTNAGTCCCTGAGCCCCGCGAGCCCGGGCCGCACACGCGACATGGGCGGCANNCCAGGG

mix

GGTACCGAGCTCGATCCACTAGTAACGCCGCCAGTGTGCTCTAAACGCCACCTGCAGGGCTACTG
CTGCTCCGGCCACTGCCTGAGACTCACCTTGCTGGAACGTGAGCCTCGGCTTCTGTCATCTCTGTG
GCCTCTGTCGCTTCTGTCGCTGTCCCCCTTNAGTCCCTGAGCCCCGCGAGCCCGGGCCGCACACGC

OSA3.1

long read
TGTGCTCGGTGTGGTTATGATCCTCGTGATGCCCTCGCTCATCAAACAGCAGGTACTGAAGAATGT
CCGCATAGACCCCAGCAGCCTGTCCTTTGCAATGTGGAAGGAGATCCCTGTACCCTTCTACTTGTC
CGTCTACTTCTTCGAGGTGGTCAATCCCAGCGAGATCCTAAAGGGTGAGAA

mix T7 and OSA3.1

GGTACCGAGCTCGATCCACTAGTAACGGCCGCCAGTGTGCTCTAAACGCCACC
TGCAGGGCTACTGCTGCTCCGGCCACTGCCTGAGACTCACCTTGCTGGAACGTG
AGCCTCGGCTTCTGTCATCTCTGTGGCCTCTGTCGCTTCTGTCGCTGTCCCCCTTN
AGTCCCTGAGCCCCGCGAGCCCGGGCCGCACACGCGACATGGGCGGCACGGCC
AGGGCGCTGGGTGGGGCTGGGGCTCGTGGGCTGCTGTGCGCTGTGCT
CGGTGTGGTTATGATCCTCGTGATGCCCTCGCTCATCAAACAGCAGGTACTGA
AGAATGTCCGCATAGACCCCAGCAGCCTGTCCTTTGCAATGTGGAAGGAGATC
CCTGTACCCTTCTACTTGTCCGTCTACTTCTTCGAGGTGGTCAATCCCAGCGAG
ATCCTAAAGGGTGAGAA

Reference: Gish, Warren and David J. States (1993). Identification of protein coding regions by database similarity search. Nature Genetics 3:266-72. Altschul, Stephen F., Warren Gish, Webb Miller, Eugene W. Myers, and David J. Lipman (1990). Basic local alignment search tool. J. Mol. Biol. 215:403-410.

Notice: statistical significance is estimated under the assumption that the equivalent of one reading frame in the query sequence codes for protein and that significant alignments will involve only coding reading frames.

Query= TITLE phasr3.seq (447 letters)

Translating both strands of query sequence in all 6 reading frames

Database: Non-redundant PDP+SwissProt+PIR+SPupdate+GenPept+GPupdate,

EDT

96,634 sequences; 27,090,059 total letters.

Searching......done

Smallest Poisson Reading High Probability Sequences producing High-scoring Segment Pairs: Frame Score P(N) LYSOSOME MEMBRANE PROTEIN II (L... +2 114 1.1e-08 1 sp|P27615|LIM2 RAT 109 6.3e-08 pir|JQ1523|JQ1523 lysosomal membrane 85K sialogly... +2 1 2.4e-06 HOMEOBOX PROTEIN HOX-2.6. >pir | ... -2 61 2 sp[P10284]HM26 MOUSE PLATELET GLYCOPROTEIN IV (GPIV) ... +2 94 1.1e-05 1 sp|P16671|CD36 HUMAN 1.1e-05 +2 94 qp|L06850|HUMCD36B 1 antigen CD36 [Homo sapiens] 1 FAT gene product [Rattus norveg... +2 92 2.3e-05 qp|L19658|RATFAT 1 1 3.8e-05 mucin - human (fragment) | 0.0 ... -1 60 2 pir(A43932(A43932 homeotic protein Hox B4 - human... -2 57 4.0e-05 pir|B60492|B60492 sp|Q01200|PRIA_LENED PRIA PROTEIN. >pir|S23106|S2310... -1 62 5.6e-05 pir|S12968|S12968 Acrosin, sperm - Pig #EC-number... -2 59 6.7e-05 88 9.0e-05 1 gp | L23108 | MUSCDANTI 1 CD36 antigen [Mus musculus] mucin - human (fragment) | 0.0 ... -1 9.2e-05 2 pir|A45106|A45106 60 Cvx peptide - Rat | 0.0 0.0 0.0... -3 2 57 0.00012 pir|S31976|S31976 2 Mox-2 [Mus musculus] -3 57 0.00012 qp|Z16406|MOX2A 1 2 Gax peptide [Rattus norvegicus] -3 57 0.00012 gp|Z17223|RNGAXMR 1 EXTENSIN PRECURSOR (CELL WALL H... -2 56 0.00024 2 splP13983 | EXTN TOBAC 1 repetitive protein antigen 69/7... -2 81 0.00035 pir|G60110|G60110 Mouse epidermal profilaggrin mR... -3 71 0.0044 1 gp | M14721 | MUSFGNAA_1 S59/4 homeotic protein - fruit ... -3 76 0.0080 pir|B36664|B36664

Plus Strand HSPs:

Score = 114 (55.2 bits), Expect = 1.1e-08, P = 1.1e-08 dentities = 22/64 (34%), Positives = 36/64 (56%), Frame = +2

Hery: 254 LLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNPSEI 433

LL + +++ V + Q + KN+ + + F W++ P+P Y+ YFF V NP EI

Sbjct: 16 LLVTSVTLLVARVFQKAVDQTIEKNMVLQNGTKVFDSWEKPPLPVYIQFYFFNVTNPEEI 75

Query: 434 LKGE 445 L+GE Sbjct: 76 LQGE 79

Plus Strand HSPs:

Score = 109 (52.8 bits), Expect = 6.3e-08, P = 6.3e-08Identities = 21/64 (32%), Positives = 35/64 (54%), Frame = +2

Query: 254 LLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNPSEI 433 LL + +++ V + Q + K + + + + F W++ P+P Y YFF V NP EI Sbjct: 16 LLVTSVTLLVARVFQKAVDQSIEKKIVLRNGTEAFDSWEKPPLPVYTQFYFFNVTNPEEI 75

Query: 434 LKGE 445 L+GE Sbjct: 76 LRGE 79

>sp|P10284|HM26_MOUSE HOMEOBOX PROTEIN HOX-2.6. >pir|A31757|A31757 homeotic protein Hox 2.6 - mouse | 0.0 0.0 0.0 0.0 0.0 >gp|M36654|MUSHOX26_1 Mouse homeo box 2.6 (Hox-2.6) mRNA, complete cds. [Mus musculus] Length = 250

Minus Strand HSPs:

Score = 61 (29.7 bits), Expect = 0.72, P = 0.52Identities = 13/41 (31%), Positives = 19/41 (46%), Frame = -2

Query: 251 PRRPAPPPPSALAVPPMSRVRPGLAGLRDSRGTATEATEAT 129
P P PPPP + P + V+P L G +EA ++
Sbjct: 75 PPPPPPPPPPPPPGLSPRAPVQPTAGALLPEPGQRSEAVSSS 115

Score = 60 (29.3 bits), Expect = 2.4e-06, Poisson P(2) = 2.4e-06 Identities = 13/25 (52%), Positives = 13/25 (52%), Frame = -2

Query: 278 PHRAQRTAAPRRPAPPPPSALAVPP 204
P QR AA R P PPPP PP
Sbjct: 59 PCTVQRYAACRDPGPPPPPPPPPP 83

>sp|P16671|CD36_HUMAN PLATELET GLYCOPROTEIN IV (GPIV) (GPIIIB) (CD36_ANTIGEN).
>pir|A30989|A30989 CD36_protein - human | 0.0 0.0 0.0 0.0 0.0
>gp|M24795|HUMANTCD36_1 Human CD36_antigen mRNA, complete cds.

[Homo sapiens] >gp|M98398|HUMCD3613_1 antigen CD36 [Homo sapiens] >gp|M98399|HUMCD3621_1 antigen CD36 [Homo sapiens] Length = 472

Plus Strand HSPs:

Score = 94 (45.5 bits), Expect = 1.1e-05, P = 1.1e-05 Identities = 18/64 (28%), Positives = 36/64 (56%), Frame = +2

Query: 245 VVGLLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNP 424 V+G + AV G +++ V LI++ + K V ++ +++F W + Y + F+V NP

Sbjct: 14 VIGAVLAVFGGILMPVGDLLIQKTIKKQVVLEEGTIAFKNWVKTGTEVYRQFWIFDVQNP 73

Query: 425 SEIL 436 E++ Sbjct: 74 QEVM 77

>gp|L06850|HUMCD36B_1 antigen CD36 [Homo sapiens] Length = 472

Plus Strand HSPs:

Score = 94 (45.5 bits), Expect = 1.1e-05, P = 1.1e-05 Identities = 18/64 (28%), Positives = 36/64 (56%), Frame = +2

Query: 245 VVGLLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNP 424
V+G + AV G +++ V LI++ + K V ++ +++F W + Y + F+V NP

Sbjct: 14 VIGAVLAVFGGILMPVGDLLIQKTIKKQVVLEEGTIAFKNWVKTGTEVYRQFWIFDVQNP 73

Query: 425 SEIL 436 E++ Sbjct: 74 QEVM 77

>gp|L19658|RATFAT_1 FAT gene product [Rattus norvegicus]
Length = 472

Plus Strand HSPs:

Score = 92 (44.5 bits), Expect = 2.3e-05, P = 2.3e-05Identities = 18/65 (27%), Positives = 36/65 (55%), Frame = +2

Query: 245 VVGLLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNP 424 V+G + AV G +++ V LI++ + + V ++ +++F W + Y + F+V NP

Sbjct: 14 VIGAVLAVFGGILMPVGDLLIEKTIKREVVLEEGTIAFKNWVKTGTTVYRQFWVFDVQNP 73

Query: 425 SEILK 439 E+ K Sbjct: 74 EEVAK 78

>pir|A43932|A43932 mucin - human (fragment) | 0.0 0.0 0.0 0.0 0.0 |
>gp|M74027|HUMMUC2A_1 mucin [Homo sapiens]
Length = 573

Minus Strand HSPs:

Score = 60 (29.1 bits), Expect = 1.4, P = 0.74Identities = 12/21 (57%), Positives = 14/21 (66%), Frame = -1

Query: 279 TTPSTAHSSPTTPSPTATQRP 217

TTPS ++ TTPSPT T P

Sbjct: 377 TTPSPPPTTMTTPSPTTTPSP 397

Score = 58 (28.1 bits), Expect = 3.8e-05, Poisson P(2) = 3.8e-05 Identities = 12/20 (60%), Positives = 14/20 (70%), Frame = -1

Query: 285 IITTPSTAHSSPTTPSPTAT 226

I TTPS ++ TTPSPT T

Sbjct: 343 ITTTPSPPTTTMTTPSPTTT 362

>pir|B60492|B60492 homeotic protein Hox B4 - human | 0.0 0.0 0.0 0.0 0.0 Length = 251

Minus Strand HSPs:

Score = 57 (27.8 bits), Expect = 2.9, P = 0.95Identities = 12/21 (57%), Positives = 12/21 (57%), Frame = -2

Query: 266 QRTAAPRRPAPPPPSALAVPP 204

QR AA R P PPPP PP

Sbjct: 63 QRYAACRDPGPPPPPPPPPPPPP 83

Score = 56 (27.3 bits), Expect = 4.0e-05, Poisson P(2) = 4.0e-05 Identities = 11/20 (55%), Positives = 12/20 (60%), Frame = -2

Query: 254 APRRPAPPPPSALAVPPMSR 195

+PR PAPPP AL P R

Sbjct: 90 SPRAPAPPPAGALLPEPGQR 109

>sp[Q01200]PRIA_LENED PRIA PROTEIN. >pir|S23106|S23106 priA protein - Shiitake mushroom | 0.0 0.0 0.0 0.0 0.0 >gp[X60956|LEPRIA_1 priA gene product [Lentinus edodes]
Length = 258

Minus Strand HSPs:

Score = 62 (30.0 bits), Expect = 0.61, P = 0.46Identities = 13/31 (41%), Positives = 18/31 (58%), Frame = -1

Query: 318 TCCLMSEGITRIITTPSTAHSSPTTPSPTAT 226

TCCL + TPS+AH + T SP++T

Sbjct: 90 TCCLPKWPTSTPTPTPSSAHHTSTHTSPSST 120

Score = 56 (27.1 bits), Expect = 5.6e-05, Poisson P(2) = 5.6e-05 Identities = 13/33 (39%), Positives = 16/33 (48%), Frame = -1

Query: 276 TPSTAHSSPTTPSPTATQRPGRAAHVACAARAR 178

TPS+ +TP P+AT G H A AR

Sbjct: 143 TPSSPSKPSSTPKPSATPNKGNGHHYKRAHVAR 175

>pir|S12968|S12968 Acrosin, sperm - Pig #EC-number 3.4.21.10 | 0.0 0.0 0.0 0.0

0.0 Length = 374

Minus Strand HSPs:

Score = 59 (28.8 bits), Expect = 1.6, P = 0.79Identities = 14/48 (29%), Positives = 24/48 (50%), Frame = -2

Query: 251 PRRPAPPPPSALAVPPMSRVRPGLAGLRDSRGTATEATEATEATEAEA 108 P++ + PP AL+ +++

LG SG+ Sbjct: 326 PQQVSAKPPQALSFAKLQQLIEALKGTAFSSGRSYYETETTDLQELPA 373

Score = 56 (27.3 bits), Expect = 6.7e-05, Poisson P(2) = 6.7e-05Identities = 11/21 (52%), Positives = 12/21 (57%), Frame = -2

Query: 266 ORTAAPRRPAPPPPSALAVPP 204

Q + PR PAPPPP PP Sbjct: 294 QPGSRPRPPAPPPPPPPPPPPPPP 314

>gp|L23108|MUSCDANTI_1 CD36 antigen [Mus musculus] Length = 473

-Plus Strand HSPs:

Score = 88 (42.6 bits), Expect = 9.0e-05, P = 9.0e-05Identities = 17/65 (26%), Positives = 35/65 (53%), Frame = +2

Query: 245 VVGLLCAVLGVVMILVMPSLIKQQVLKNVRIDPSSLSFAMWKEIPVPFYLSVYFFEVVNP 424

LI++ + + V ++ + +F W +

Sbjct: 15 VIGAVLAVFGGILMPVGDMLIEKTIKREVVLEEGTTAFKNWVKTGTTVYRQFWIFDVQNP 74

Query: 425 SEILK 439

++ K

The Arthur Standard Standard

Sbjct: 75 DDVAK 79